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**BLAST 2 sequences** 

<u>BLAST</u>

**Example** 

*Help* 

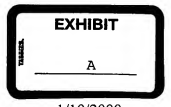
## **BLAST 2 SEQUENCES**

This tool produces the alignment of two given sequences using <u>BLAST</u> engine for local alignment. Currently only blastn and blastp programs are available. Using sequences > 150 Kb is not recommended.

Reference: Tatiana A. Tatusova, Thomas L. Madden (1999), "Blast 2 sequences - a new tool for comparing protein and nucleotide sequences", FEMS Microbiol Lett. 174:247-250

Program   blastp   Matrix   0 BLOSUM62	
Parameters used in BLASTN program only:	
Reward for a match: Penalty for a mismatch:	
Open gap 11 and extension gap 1 penalties	,
gap x_dropoff 50 expect 10000 word size Filter Filter	
Sequence 1 Enter accession or GI	Light Chain CDRI
or sequence in FASTA format from:	,
KTSQDINKYMA	X
	Marine al.G
Sequence 2 Enter accession or GI	•
or sequence in FASTA format from:	
RSSQSLAKSYGNTYLS  Align Clear Input	lesidues 44-59 of SEQ ID NO:12
Comments and suggetstions to: blast-help@ncbi.nlm.nih.gov	

Credits to: <u>Tatiana Tatusov</u> and Tom Madden



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**BLAST 2 sequences** 

<u>BLAST</u>

<u>Help</u>

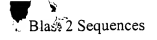
BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.0.9 [May-07-1999]

Matrix 0 BLOSUM62 gap open: 11 gap extension: 1 x\_dropoff: 50 expect: 1000c wordsize: 2 Filter Align

Sequence 1 lcl|seq\_1 Length 11

Sequence 2 lcl|seq\_2 Length 16

No siginificant similarity was found



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**BLAST 2 sequences** 

<u>BLAST</u>

**Example** 

<u>Help</u>

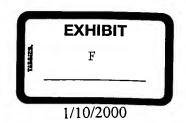
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Program blastp Matrix 0 BLOSUM62	***************************************
Parameters used in BLASTN program only:	_
Reward for a match: Penalty for a mismatch:	
Open gap 11 and extension gap 1 penalties	
gap x_dropoff 50 expect 1000c word size 2 Filter Filter	
Sequence 1 Enter accession or GI Heavy Chair CDR	3
or sequence in FASTA format from:to:	,
EGYYGNYGVYAMDY Muint	21.6
EGYYGNYGVYAMDY  Muinl	
For a second sec	
Sequence 2 Enter accession or GI	
or sequence in FASTA format from:to:	
GGYDGWDYAIDY Pesidues II	8-129
SEQ ID NO	; 15
Align Clear Input	
	and the second s
Comments and suggetstions to: <u>blast-help@ncbi.nlm.nih.gov</u>	

Credits to: Tatiana Tatusov and Tom Madden





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**BLAST 2 sequences** 

<u>BLAST</u>

<u>Help</u>

BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.0.9 [May-07-1999]

Matrix 0 BLOSUM62 

gap open: 11 gap extension: 1

x\_dropoff: 50 expect: 1000c wordsize: 2 Filter Align;

Sequence 1 lcl|seq\_1 Length 14

Sequence 2 lcl|seq 2 Length 12

No siginificant similarity was found

1 4

<u>NCBI</u> Entrez. **BLAST 2 sequences** 

<u>BLAST</u>

**Example** 

<u>Help</u>

## **BLAST 2 SEQUENCES**

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Reference: Tatiana A. Tatusova, Thomas L. Madden (1999), "Blast 2 sequences - a new tool for comparing protein and nucleotide sequences", FEMS Microbiol Lett. 174:247-250

Program   blastp   Matrix   U BLOSUM62	
Parameters used in BLASTN program only:	<del>-</del>
Reward for a match: Penalty for a mismatch:	
Open gap and extension gap penalties	
gap x_dropoff 50 expect 1000c word size 2 Filter Filter	
Sequence 1 Enter accession or GI	Light Chain CDR3
or sequence in FASTA format from: to:	/
LQYDNLWT	Murine 21.6
\$	Murine 21.6
Sequence 2 Enter accession or GI	
or sequence in FASTA format from:to:	· •••
LQGTHQPYT	Residues 114-122 of SEO ID NO:12
	SEO ID NO: 12
Align Clear Input	
Comments and suggetstions to: <u>blast-help@ncbi.nlm.nih.gov</u>	

Credits to: <u>Tatiana Tatusov</u> and <u>Tom Madden</u>



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**BLAST 2 sequences** 

<u>BLAST</u>

<u>Help</u>

BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.0.9 [May-07-1999]

Matrix 0 BLOSUM62 gap open: 11 gap extension: 1

x\_dropoff: 50 expect: 1000c wordsize: 2 Filter Allign

Sequence 1 lcl|seq\_1 Length 8

Sequence 2 |cl|seq\_2 Length 9

No siginificant similarity was found